Attorney Docket No.:

TLHR-0005 (124187.00009.US2)

AMENDMENT TO THE CLAIMS

What is Claimed is:

1. (Currently Amended) A method of diagnosing or monitoring a lysosomal storage disorder in a patient, comprising:

obtaining a first sample from the patient; and

measuring a first level of at least a first saposin in the first sample obtained from the patient;

comparing the first level to a baseline level, wherein the baseline level is the level of at least the first saposin as determined in a control population of patients unaffected by the lysosomal storage disorder; and

determining a presence or extent of a lysosomal storage disorder when the first level is similar or different than the 95th percentile of the baseline level of at least the first saposins in the control population;

wherein,

- (i) the first level is similar or different from the baseline level;
- (i) the similarity of the first level compared to the baseline level is an indicator of absence of the lysosomal storage disorder in the patient;
- (ii) the <u>difference of the</u> first level <u>compared to the baseline level</u> is an indicator of presence or extent of the lysosomal storage disorder in the patient;
 - (iii) the first saposins comprises saposin A, saposin B, saposin C, saposin D, presaposin, mRNA encoding presaposin, or a combination thereof; and
 - (iv) the first sample is a plasma, serum, whole blood, urine, or amniotic fluid sample.
- 2. (Canceled) The method of claim 1, wherein the first sample is a plasma sample.
- 3. (Canceled) The method of claim 1, wherein the first sample is a whole blood sample.
- 4. (Currently Amended) The method of claim 1, <u>further comprising indicating a presence of the lysosomal disorder in the patient when the first level exceeds the baseline level.</u> wherein a presence of the lysosomal disorder in the patient, is indicated by the first level exceeding the baseline level.

Attorney Docket No.: TLHR-0005 (124187.00009.US2)

5. (Currently Amended) The method of claim 1, further comprising:

measuring a second level of the <u>a second</u> saposin in a second sample from the patient, <u>wherein the first saposin and second saposin are the same, and</u> the first and second samples being <u>are</u> obtained at different times; and

comparing the first level and the second level in the samples to monitor progression of the disease,

determining a presence or extent of a lysosomal storage disorder when the second level is similar or different than the 95th percentile of the baseline level of at least the two saposins in the control population;

wherein,

- (i) the second saposin comprises saposin A, saposin B, saposin C, saposin D prosaposin, mRNA encoding prosaposin, or a combination thereof;
- (ii) the comparison of the first level and the second <u>level</u> is an indicator of the progression of the disease in the patient; and
- (iii) the second sample is a plasma, serum, whole blood, urine, or amniotic fluid sample.
- 6. (Currently Amended) The method of claim 1, wherein the patient further comprising selecting the patient that is undergoing treatment for the lysosomal storage disorder.
- 7. (Canceled) The method of claim 4, wherein the first level is greater than the 95th percentile of the baseline level in the control population.
- 8. (Currently Amended) The method of claim 1, wherein the patient further comprising selecting the patient that is not known to have a lysosomal storage disorder before the measuring step.
- 9. (Currently Amended) The method of claim 1, wherein the patient further comprising selecting the patient that is an infant less than one year old.
- 10. (Currently Amended) The method of claim 1, <u>further comprising selecting the patient</u> that wherein the patient is a fetus and the sample is a fetal blood sample.

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11. (Previously Presented) The method of claim 5, wherein a change in the first level of the saposin indicates progression or regression of the disorder in the patient that is known to have a lysosomal storage disorder.

- 12. (Previously Presented) The method of claim 5, wherein a change in the first level of the saposin indicates a response to treatment of the lysosomal storage disorder in the patient that being treated for the lysosomal storage disorder.
- 13. (Canceled) The method of claim 5, wherein the first sapesin or second sapesin is selected from the group consisting of sapesin A, sapesin B, sapesin C, and sapesin D.
- 14. (Canceled) The method of claim 1, wherein the sapesin is selected from the group consisting of sapesin A, supesin C, or sapesin D.
- 15. (Previously Presented) The method of claim 1, wherein the measuring step comprises detecting binding between a saposin polypeptide and an antibody.
- 16. (Original) The method of claim 15, wherein the antibody is a monoclonal antibody.
- 17. (Original) The method of claim 15, wherein the antibody is immobilized to a solid phase.
- 18. (Currently Amended) The method of claim 1, wherein the lysosomal storage disorder is selected from the group consisting of cystinosis, Fabry's disease, Niemann-Pick disease, Pompe's disease, Wolman disease, and subset thereof.
- 19. (Original) The method of claim 1, further comprising informing the patient or a parent or guardian thereof of the presence of the lysosomal storage disorder.
- 20. (Previously Presented) The method of claim 1, further comprising determining a treatment program based on the measurement of the first level of the first saposin.
- 21. (Withdrawn) A method of diagnosing or monitoring a lysosomal storage disorder in a patient, comprising: measuring the level of a-glucosidase in a tissue sample from a patient, wherein the level is an indicator of the presence or extent of the disorder in the patient.
- 22. (Withdrawn) The method of claim 21, wherein the sample is a plasma sample.

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PATENT

Attorney Docket No.: TLHR-0005 (124187.00009.US2)

23. (Withdrawn) The method of claim 21, wherein the sample is a blood sample.

- 24. (Withdrawn) The method of claim 21, further comprising diagnosing the presence of a disorder selected from the group consisting of acid lipase disease, mannosidosis, MPSII, MPS IIIA, MSD, mucolipidosis, N-P (A/B), N-P (C), Sandhoff, SAS or TSD B1, if the measured level of a-glucosidase exceeds the mean level in a control population of individuals not having a lysosomal storage disease.
- 25. (Withdrawn) The method of claim 21, further comprising diagnosing the presence of disorder selected from the group consisting of galactosialidosis, MPS IVA and Pompe's disease if the measured level of a-glucosidase is below the mean level in a control population of individuals not having lysosomal storage disease.
- 26. (Withdrawn) A method of diagnosing a lysosomal storage disorder comprising measuring a level of a saposin in a tissue sample from the patient; measuring a level of LAMP-1 or LAMP-2 in a second tissue sample from the patient; measuring a level of a glucosidase in a third tissue sample from the patient; wherein an increased level of saposin and/or LAMP-1 or LAMP-2, and/or an increased or decreased level of a-glucosidase in the sample relative to respective mean levels in a control population is an indicator of presence or extent of the disorder in the patient.
- 27. (Withdrawn) A method of diagnosing Pompe's disease in a patient, comprising measuring a level of a saposin in a tissue sample from the patient: measuring the level of a-glucosidase in a second tissue sample from the patient; wherein the presence of an increased level of the saposin and a decreased level of the a-glucosidase relative to mean levels of the saposin and a-glucosidase in a control population of individuals not having a lysosomal storage disorder indicates Pompe's disease or susceptibility thereto.
- 28. (Withdrawn) A method of screening patients for presence of lysosomal storage disorder, comprising: measuring the level of a LAMP-1 polypeptide in a sample from the patient: measuring the level of a saposin peptide in the sample, the presence of an increased level of LAMP-1 or saposin or both relative to mean levels in a control population, indicating susceptibility to a lysosomal disorder.

Attorney Docket No.: TLHR-0005 (124187.00009.US2)

29. (Withdrawn) A diagnostic kit comprising: a first reagent that binds to a LAMP; a second reagent that binds to a saposin.

- 30. (Withdrawn) The diagnostic kit of claim 29, further comprising a third reagent that binds to a glucosidase.
- 31. (Withdrawn) The diagnostic kit of claim 30, wherein the first, second and third reagents are antibodies.
- 32. (Withdrawn) In a method of screening a patient for presence or susceptibility to disease, comprising performing a plurality of diagnostic tests on a tissue sample from the patient for a plurality of diseases, the improvement wherein one of the diagnostic tests comprises measuring the level of a saposin.
- 33. (Withdrawn) In the method of claim 32, the further improvement wherein a second of the diagnostic tests comprising measuring the level of LAMP-1 in the tissue sample from the patient.
- 34. (Withdrawn) In the method of claim 33, the further improvement wherein a third of the diagnostic tests comprises measuring the level of a-glucosidase in the tissue sample from the patient.
- 35. (Withdrawn) In the method of claim 32, the further improvement wherein a fourth of the diagnostic test comprises analysing a nucleic acid encoding an enzyme associated with a lysosomal storage disorder for a polymorphic form correlated with the disorder.
- 36. (Previously Presented) A method of monitoring treatment of a lysosomal storage disease in a patient, comprising:

determining a pre-treatment baseline level of a saposin in a sample from the patient with a lysosomal storage disorder before treatment with an agent;

determining a post-treatment baseline level of the saposin in a sample from the patient with the lysosomal storage disorder after treatment with the agent; and

comparing the pre-treatment baseline level of the with the post-treatment baseline level of the saposin, wherein

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- (i) the sample is a plasma, serum, whole blood, urine, amniotic fluid sample, or a mixture of;
- (ii) saposin is selected from the group consisting of saposin A, saposin B, saposin C, saposin D, prosaposin, mRNA encoding presaposin, and a combination thereof; and
- (iii) a reduction in the post-treatment baseline level relative to the pre-treatment baseline level indicates a positive treatment outcome.
- 37. (Withdrawn) A method of monitoring treatment of acid lipase disease, mannosidosis, MPSII, MPS IIIA, MSD, mucolipidosis, N-P (A/B), N-P (C), Sandhoff, SAS or TSD B1, comprising: determining a baseline level of a glucosidase in a tissue sample from the patient with the disorder before treatment with an agent; comparing a level of the a glucosidase in a tissue sample from the patient with the disorder after treatment with the agent with the baseline level; wherein a decrease relative to the baseline indicates a positive treatment outcome.
- 38. (Withdrawn) A method of monitoring a patient with Pompe's disease, comprising: determining a baseline level of a glucosidase in a tissue sample from the patient with the disorder before treatment with the agent; comparing a level of the a-glucosidase in a tissue sample from the patient after treatment with the agent with the baseline level; wherein an increase relative to the baseline indicates a positive treatment outcome.
- 39. (New) A method of diagnosing or monitoring a lysosomal storage disorder in a patient, comprising:

 obtaining a first sample from the patient;
 measuring a first level of a saposin in the first sample obtained from the patient:
 comparing the first level to a baseline level, wherein the baseline level is the level of the
saposin as determined in a control population of patients unaffected by the lysosomal
storage disorder;
determining a presence or extent of a lysosomal storage disorder when the first level is
similar or different than the 95th percentile of the baseline level of at least the two
saposins in the control population:

wherein,

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Fax:9727442909

Dec 2 2005 16:10

P. 10

PATENT

Attorney Docket No.: TLHR-0005 (124187.00009.US2)

	(i) the similarity of the first level compared to the baseline level is an indicator of
absence of the	lysosomal storage disorder in the patient;
	(ii) the difference of the first level compared to the baseline level is an indicator
of presence or	extent of the lysosomal storage disorder in the patient;
	(iii) the saposin comprises saposin A, saposin B, saposin C, saposin D;
	(iv) the first sample is plasma; and
	(v) the baseline level and the first level are about equal to a percent elevation level
	for the lysosomal storage disorder listed in Table 2.